



TITLE:

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ANISEED 2017: extending the integrated ascidian database to the exploration and evolutionary comparison of genome-scale datasets

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ABSTRACT

ANISEED (www.aniseed.cnrs.fr) is the main model organism database for tunicates, the sister-group of

vertebrates. This release gives access to annotated genomes, gene expression patterns, and anatomical descriptions for nine ascidian species. It pro-

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vides increased integration with external molecular and taxonomy databases, better support for epigenomics datasets, in particular RNA-seq, ChIP-seq and SELEX-seq, and features novel interactive interfaces for existing and novel datatypes. In particular, the cross-species navigation and comparison is enhanced through a novel taxonomy section describing each represented species and through the implementation of interactive phylogenetic gene trees for 60% of tunicate genes. The gene expression section displays the results of RNA-seq experiments for the three major model species of solitary ascidians. Gene expression is controlled by the binding of transcription factors to *cis*-regulatory sequences. A high-resolution description of the DNA-binding specificity for 131 *Ciona robusta* (formerly *C. intestinalis* type A) transcription factors by SELEX-seq is provided and used to map candidate binding sites across the *Ciona robusta* and *Phallusia mamillata* genomes. Finally, use of a WashU Epigenome browser enhances genome navigation, while a Genomicus server was set up to explore microsynteny relationships within tunicates and with vertebrates, *Amphioxus*, echinoderms and hemichordates.

INTRODUCTION

The tunicate clade of marine invertebrate deuterostomes includes the sessile ascidians and the pelagic appendicularians and thaliaceans (1,2). A major interest of the scientific community for tunicates stems from the phylogenetic position of this chordate group, now considered to be the long-diverged sister group of vertebrates (3). Solitary ascidians and all appendicularians reproduce sexually only. By contrast, colonial ascidians and all thaliaceans reproduce both sexually and asexually by budding (1).

Ascidians, a paraphyletic group (4), are the largest and most studied group of tunicates. Major solitary ascidian model species, studied for their embryogenesis, aging and regeneration properties are *Ciona robusta* (formerly *Ciona intestinalis* type A), *Ciona intestinalis* (formerly *Ciona intestinalis* type B), *Ciona savignyi*, *Phallusia mamillata*, *Halocynthia roretzi* and *Molgula oculata*. The colonial ascidian *Botryllus schlosseri* is well known for studies on whole-body regeneration and stem cells (5), and on self/non-self recognition (6). Comparison of ascidian and vertebrate strategies can provide cues on the origin of vertebrate novelties such as the neural crest (7) or the second heart field (8). Their rapid molecular evolution is also informative on the plasticity of the chordate program of gene regulation (8–11) or of histocompatibility (6). In addition to this basic science focus, many solitary and colonial ascidian species are invasive, major contributors to bio fouling and, as such, pests for the aquaculture industry (12,13).

Over the past 20 years, an extensive functional genomics toolbox (14) has been developed to make best use of the simplicity of the embryo and genome (15) of *Ciona intestinalis* and *robusta*. This tool has now been extended/adapted to other tunicate species (16,17). Interference with gene func-

tion, initially relying on morpholino oligonucleotides, now includes the use of TALENs (18) and CRISPR/cas9 strategies (19,20).

ANISEED - Ascidian Network for In Situ Expression and Embryological Data—is the major database system in the Tunicate community, hosting and giving access to genomic, genetic and anatomical data on ascidian development. First published in 2010 (21), the database hosts and gives access to genomic, genetic and anatomical data on tunicates. Its refactoring using the Chado schema philosophy and its extension to 11 ascidian species was completed in 2015 (22). Since then, tunicate laboratories have rapidly embraced genomics, including ChIP-seq (23,24), RNA-seq of whole embryos (24) or FACS-sorted single cells (25), bisulphite sequencing (26), SELEX-seq and TSS-seq (27). The aim of the ANISEED 2017 update is to accompany the entry of tunicate biology into the genome-scale era through three types of improvements: extension/refactoring of mining and display tools, extension of pre-existing datasets and development of modules to host and give access to novel types of genomics data.

NEW DATA MINING/DISPLAY TOOLS

Taxonomy pages

A taxonomy page accessible from the home page was introduced in ANISEED 2017 to place the covered species into their phylogenetic context and facilitate the interpretation of comparative studies. The page shows a cladogram of the twelve species currently described at the anatomical or genetic level in the database. Each node gives access to a page describing the corresponding species. Following the novel taxonomy (28) and a community decision at the 2017 international Tunicate Meeting, *Ciona intestinalis* type A has been renamed *Ciona robusta* throughout the database; while *Ciona intestinalis* type B is now referred to as *Ciona intestinalis*. No molecular data are currently associated to this latter species in ANISEED, as its genome draft is of insufficient quality to build gene models.

Each species page briefly describes the species, its distribution and its ecology, lists the number of ANISEED entries and provides a link to the corresponding ANISEED genome browser. This page also links to other tunicate molecular databases (DBTGR (29), CiPro (30), Ghost (31) and FABA (32) for *Ciona robusta*), to the CRISPOR CRISPR/Cas9 guide RNA design tool for ascidian species (33), as well as to major ecological databases, including the Ascidiacea World Database from the World Registry of Marine species (WoRMS) (34), the Global Biodiversity Information Facility (GBIF) (35), or the Global Invasive Species Database (<http://www.invasivespecies.net/>). Finally, for practical purposes, the page lists for each species the animal providers used in the scientific community.

Interactive gene phylogenies. The diversity of ascidian species with an assembled genome provides a framework to study the evolution of developmental mechanisms. To identify orthologous genes within tunicates and with chordates, ANISEED 2015 used an orthology pipeline based on the OrthoMCL software (36) and listed for each gene the cluster of homologous genes it was associated to. In many cases,

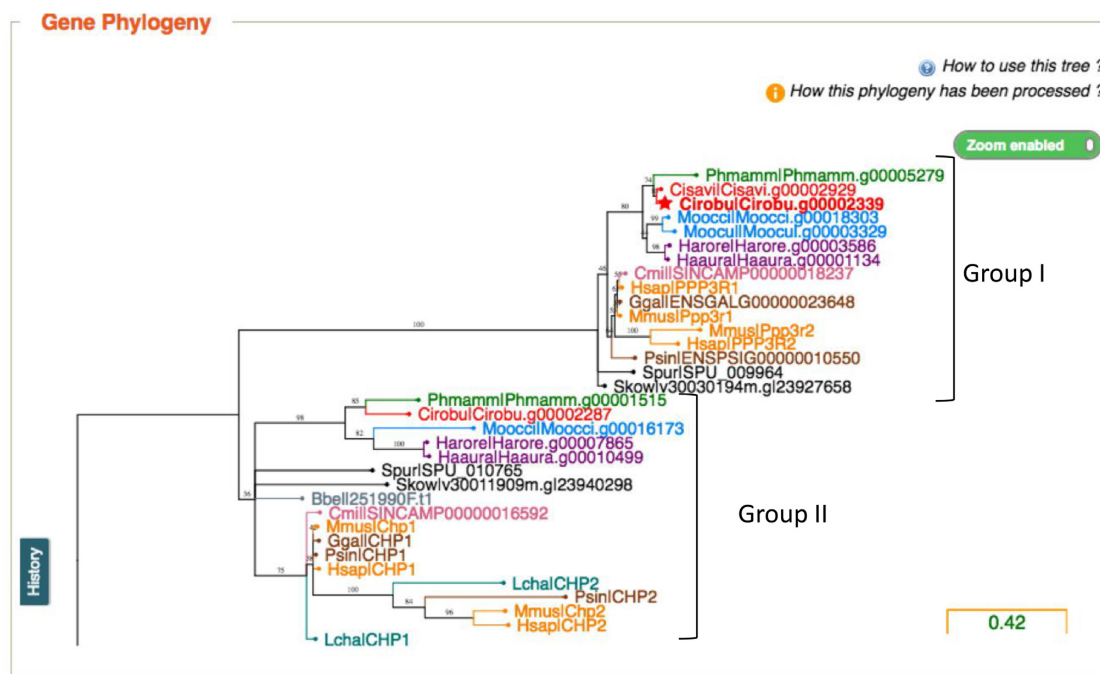


Figure 1. Interactive gene Phylogenies. Screenshot from a part of the phylogenetic gene tree including genes *Cirobu.g00002339* and *Cirobu.g00002287*. Species are color-coded and the tree can be compressed or extended lengthwise and zoomed in and out. Note the difference between the two groups of genes. In group I, the gene phylogeny follows the species phylogeny, and tunicate genes are considered orthologous to the vertebrate genes of the group. In group II by contrast, the gene phylogeny does not follow the species phylogeny, and we consider that the tunicate genes have no unequivocal vertebrate orthologs.

clusters were large, and in the absence of phylogenetic trees, the system did not allow the identification of accurate 1-to-1 orthology relationships within the list.

We thus refactored the ANISEED 2017 orthology pipeline. We first used SiLiX (37) to build clusters of homologous genes from 18 species (6 vertebrates, 9 ascidians, 1 lancelet and 2 non-chordate deuterostome outgroups). A phylogenetic tree was then inferred for each cluster using RaxML (38) under the LG+Γ4+F evolution model (See supplementary methods). Trees for approximately 60% of the coding gene models of each ascidian species are displayed interactively on the corresponding gene card using PhyloCanvas (<http://phylocanvas.org>) (Figure 1). This procedure identifies unequivocal vertebrate orthologs for ~30% of ascidian genes (average number of human orthologs/*C. robusta* gene: 2.8). A similar frequency of tunicate genes with clear human orthologs is also observed in ENSEMBL 90 (28% of *C. robusta* genes with high-confidence human orthologues). While this situation may reflect the large genetic distance between tunicates and vertebrates, we noticed that orthology relationships could also be confounded by the position the non-olfactorian outgroups (Figure 1). Approximately 25% of ascidian genes without clear vertebrate orthologs have at least one ortholog in an ascidian of a different order (Supplementary Table S1).

New genome browsers

ANISEED 2015 featured Gbrowse 2.0 (39) genome browsers, which showed limitations as the size of the dataset

increased. The WashU Epigenome browser (40) was selected as a replacement and adapted with the help of its developer. It offers increased speed and the ability to represent genome-wide chromatin interaction data (41). Within or between genome comparisons are facilitated by the display in a unique window of concatenations of sets of gene loci within a species, or between species (e.g. orthologous loci to explore synteny relationships).

Tracks presented are grouped into three categories. ‘Annotation tracks’ provide general background information (gene models, repeated elements, inter-species genome sequence comparison, genome alignment, operons, *cis*-regulatory sequences, Ns and GC content). ‘Public track hubs’ group tracks presenting the information gathered within a specific project or using a specific genome-scale technique (RNA-seq, TSS-seq, methyl-seq, SELEX-seq, ChIP-seq: see below). For each public hub, a precise description of the content and methods is provided. Finally, ‘Custom tracks’ can be uploaded as individual tracks or hubs. Each session can be saved and shared with other scientists. A useful wiki handbook is provided by the developer (wiki.wubrowse.org).

The WashU and Gbrowse 2.0 browsers will coexist until summer 2018, after which only the WashU browser will remain. During this 1-year transition period, the addition of new tracks, some of which are presented here, will be restricted to the WashU browser.

Tunicate genomes rapidly diverge at the local sequence level, as well as in their organization. To further explore the conservation of genome organization within deuterostomes, we implemented a Genomicus synteny browser (42),

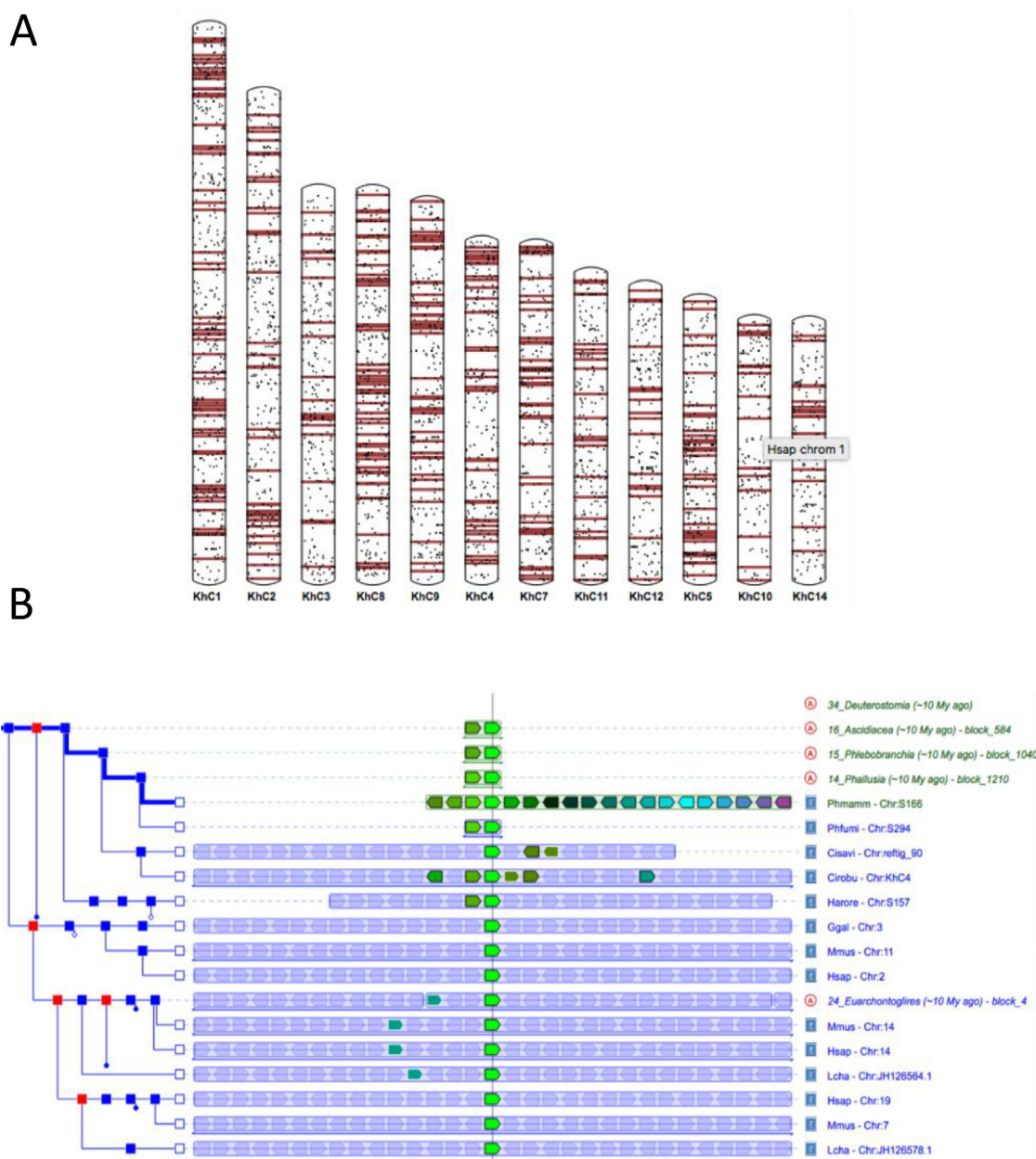


Figure 2. Genomic and the study of global and local genome rearrangements. (A) Karyotype view showing the mapping of segments of Human Chromosome 1 onto *Ciona robusta* chromosomes. (B) Phyloview of the *Phallusia mammillata* gene Phmamm.g00004580 showing rapid loss of microsynteny within tunicates and complete loss with vertebrates.

based on the refined orthology relationships described above. This tool provides powerful visualization interfaces, such as karyotype View to explore genome organization (Figure 2A) and PhyloView to display the evolution of microsynteny relationships within ascidians and with vertebrates or non-chordate deuterostomes (Figure 2B).

EXTENSION OF PREEXISTING DATA TYPES

Functional gene annotations

Functional genome annotation (transcript and gene models, orthology to other tunicates, to vertebrates and to other deuterostomes, best blastp hit to human proteins, Inter-

pro and GO annotations) was extended to all species with a genome assembly of sufficient quality to support gene model building, including three species previously only represented on the genomic browser: *Phallusia fumigata*, *Holocynthia aurantium* and *Molgula occidentalis*.

Gene models are annotated on each gene page and also provided as annotation tracks on the browsers. A link to alternative NCBI and ENSEMBL gene models is provided on each gene card. Manually-curated gene names respecting the tunicate nomenclature guidelines (43) are currently being generated by a dedicated committee and will be gradually added until the next official release. Genes can currently be searched for in ANISEED and in the WashU browser by

Table 1. Evolution of the number of entries in ANISEED between October 2010 and October 2017

Class of entries	2010		2015				2017				Phallusia mammi- lata
Species	Halocynthia roretzi	Ciona robusta	Halocynthia roretzi		Ciona robusta		Halocynthia roretzi		Ciona robusta		
	Number	Number	Number	% Increase /2010	Number	% Increase /2010	Number	% Increase /2015	Number	% Increase /2015	
Measure;	Number	Number	Number	% Increase /2010	Number	% Increase /2010	Number	% Increase /2015	Number	% Increase /2015	Number
Articles in Pubmed (query: species + gene + development)	-	214	NA	NA	280	31%	79	NA	339	21%	7
Articles in ANISEED	1	160	1	0	217	36%	4	300%	253	17%	0
Molecular Tools	0	619	0	NA	769	24%	0	NA	874	14%	0
Regulatory Regions Analyzed (literature)	0	528	0	NA	875	66%	0	NA	1102	26%	0
Spatio-temporal patterns of cis-regulatory activity (literature)	0	777	0	NA	944	21%	0	NA	1152	22%	0
Gene Expression Patterns in Mutant Embryo (literature)	0	1152	0	NA	1294	12%	51	NA	1860	44%	0
Gene Expression Patterns (literature)	28	4259	51	82%	6545	54%	154	202%	7717	18%	0
Gene Expression Patterns (large scale)	5825	21017	5825	0%	21017	0%	5825	0%	21538	2%	262
Total Gene Expression Patterns	5853	25276	5876	0%	27562	9%	5979	2%	29255	6%	262
Nb of genes described	933	4000	937	0%	4500	13%	943	1%	4750	6%	31

unique gene or gene model Ids, by name of their most related human gene, or through the ANISEED blast server, which can now blast protein or nucleotide queries against both whole genomes or transcript model sets.

The functional annotation of ascidian gene models is based on Gene Ontology (44) terms. Because of the complexity and depth of the GO ontology, containing more than 40 000 terms, genes with similar function can be annotated with distinct terms, which can confound GO term enrichment analyses (45). To overcome this issue, we used an objective framework Information Content algorithm (46, stringency parameter 0.2) to define a smaller Tunicate GO Slim of 633 terms (MF:129; BP: 374; CC: 130), which provide higher level and more consistent annotations. We mapped global GO annotations onto this GO slim and propose this annotation in the download section of the database. For *Ciona robusta*, this process led to the annotation of 69% of genes with at least one term of each MF, BP and CC category.

Manually curated data

ANISEED hosts both large-scale data, automated annotations, and manually curated data from the literature or communicated by partner laboratories. This latter category, which now forms 30% of total gene expression patterns in *Ciona robusta*, is crucial to contextualize large-scale datasets, and is the primary source of expression profiles in altered developmental conditions (gene knock-down or overexpression; 6% of *Ciona robusta* dataset). Table 1 presents the growth of major sections of ANISEED between 2010 and 2017. The majority of the new data originates from the manual curation of 33 articles describing the development of *Ciona robusta*, published between 2012 and 2016, including 19 describing neural development, a very active field of *Ciona* research. The newly curated dataset set also includes 3 *Halocynthia roretzi* articles.

NEW DATASETS

Annotated Repeated elements

Repeated elements, previously not described, were detected for all nine species, and each element was analysed and

named according to the guidelines for the nomenclature of tunicate genetic elements (43). These elements are displayed in the 'Annotation tracks' section of the WashU genomic browser.

RNA-seq

RNA-seq provides a quantitative estimation of the steady-state expression level of genes in wild-type and experimentally perturbed conditions. Gene cards in ANISEED 2017 include in a new section of the expression tab a summary of gene expression dynamics across all wild-type and mutant RNA-seq experiments analysed (Supplementary Figure S1). For each experiment two complementary but comparable normalizations are displayed: FPKM/RPKM and an 'unlogged' version of RLE (47). The former is best to compare the expression of different genes in the same experiment, while the latter is a better choice to compare the expression of a given gene across experiments. Note that 'unlogged' RLE values are not suitable for statistical analysis. The download button of the page thus returns the true logarithmic RLE value.

Two sets of data of immediate interest to the community were quantified. The first dataset is composed of unpublished stranded RNA-seq data for whole embryos of seven developmental stages ranging from egg to hatching larva in three species (*Ciona robusta*, *Phallusia mammillata* and *Halocynthia roretzi*). This dataset allows comparing the expression dynamics of orthologous genes in these three divergent ascidian species. Note that the *Ciona* data are from *Ciona intestinalis* embryos, but in the absence of a high quality *C. intestinalis* genome assembly, were mapped onto the *Ciona robusta* genome and considered to reflect the expression of the *Ciona robusta* orthologs. The second dataset (NCBI bioproject PRJDB3843) compares wild-type, Bmp-treated and Bmp-inhibited (Dorsomorphin) *Ciona robusta* late gastrula embryos. Additional datasets will be progressively integrated as they become public. Each of these datasets is also integrated as a public hub in the WashU browser.

RNA-seq usually poorly represents immature unspliced transcripts. This technique is therefore inadequate to identify Transcription Start Sites (TSS), in particular in ascidians, which make frequent use of trans-splicing (10). Re-



Figure 3. Simultaneous visualization of candidate transcription factor binding sites in two known regulatory sequences. The WashU epigenome browser allows splitting the display to show two independent loci from the same genome. Here the display shows regulatory sequences of the *Cirobu* *Otx* (*Cirobu.g00006940*; left part of the window) and *Nodal* (*Cirobu.g00010576*; right part of the window) genes, and predicted local *in silico* binding affinity for transcription factors of the ETS and GATA families, known to regulate these enhancers.

cently, high-throughput RNA-sequencing techniques were developed to produce precise and quantitative genome-wide mapping of TSS, thereby allowing a better analysis of promoters (48). The *Ciona robusta* WashU browser of ANISEED 2017 includes a public hub with TSS-seq mapping for a set of adult tissues, and whole larvae (27).

ChIP-seq

In addition to the *Ciona robusta* ChIP-chip data for 11 transcription factors previously displayed on the *Ciona robusta* genome browser (49), ChIP-seq data for three additional transcription factors at the 32-cell stage (23) are now accessible as public hubs in the *C. robusta* WashU browser. ChIP-seq for the promoter-specific H3K4me3 histone mark in *C. robusta* and *P. mammillata* are also provided to help, combined with TSS-seq data, with the identification of basal promoters.

SELEX-seq

Gene expression is controlled at the transcriptional level by non-coding *cis*-regulatory sequences that act as binding platforms for transcription factors, which recognize specific DNA sequences. The knowledge of this binding specificity is a major help in deciphering the gene regulatory networks that underlie biological function. ANISEED 2017 describes the *in vitro* DNA-binding specificity of 131 of the estimated 500 transcription factors encoded in the *Ciona robusta* genome. These DNA-binding specificities, determined by SELEX-seq (50), are accessible via a specific tab in the gene card, which describes the details of the enrichment procedure and DNA-binding specificity and the relatedness of DNA-binding specificity to mammalian orthologs

(50,51). 6-mer enrichments values and Position Weight Matrices can also be downloaded from this page. In addition, public hubs on the *Ciona robusta* and *Phallusia mammillata* WashU browsers group tracks of local predictions of binding affinity for each of these 131 *C. robusta* transcription factors and their 79 *P. mammillata* orthologs (Figure 3).

HOSTING OF PRIVATE NEW TUNICATE GENOMES

ANISEED offers to host on its genome browsers and blast servers, password-protected, private genomes sequenced by members of the community. Two genome projects are currently hosted and kept private until their publication in peer reviewed journals: the colonial stolidobranch ascidian *Botrylloides leachii* and the solitary phlebobranch *Corella inflata*.

WEBSERVICES AND DOWNLOADS

To facilitate the programmatic extraction of datasets and the development of external modules relying on ANISEED data, we developed an Application Programming Interface (API), accessible from the main menu. Currently, the main functions allow retrieving genes according to their expression patterns, articles and authors, *cis*-regulatory regions and *cis*-regulatory constructs and cDNA clones.

The download section was enriched with standardized GAF files with functional gene annotations for all species. Two functional annotations are proposed for each species, one based on the full Gene Ontology, the other on the tunicate GOslim we developed, which can also be downloaded. The gene trees are also available (.tre format). Finally, 6-mer enrichments and Position Weight Matrices for all rounds for

131 SELEXed *Ciona robusta* transcription factors are accessible.

FUTURE PERSPECTIVES

ANISEED 2017 marks a substantial improvement in the content and organization of the main tunicate model organism database. The diversity and depth of the added data attests of the dynamism of this small scientific community of less than a hundred labs worldwide.

One major evolution over the past two years was the discovery that the species initially called *Ciona intestinalis* comprises at least two independent species, *Ciona robusta* (formerly called *Ciona intestinalis* type A and found in all major oceans) and *Ciona intestinalis* (formerly called *Ciona intestinalis* type B, and mostly restricted to the North Atlantic) (28). In this version of ANISEED, we therefore reallocated all *Ciona intestinalis* data to *Ciona robusta*, which is now the most studied *Ciona* species. To facilitate interoperability and comparison between databases and to avoid confusion between species, it will be important for the *Ciona* community to also press for a reassignment of data to their correct species in generalists databases including NCBI, ENSEMBL, Uniprot or UCSC, which have so far not adapted to the new taxonomy.

We would also like to extend the database, beyond ascidians, to appendicularians and thaliaceans. While genomic information remains very partial in thaliaceans (52), high-quality genomic (53), transcriptomic (54) and functional (55) data exist for the appendicularian *Oikopleura dioica*, and we have established contact with the scientific community working on these animals. Computing accurate orthology relationships for most genes of these fast evolving taxa may, however, prove particularly challenging, and may require further optimization of our procedures.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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